



(1) Publication number: 0 510 907 A2

(12)

EUROPEAN PATENT APPLICATION

(21) Application number: 92303533.1

(22) Date of filing: 21.04.92

(f) Int. Cl.⁵: **C12N 15/55**, C12N 9/14, A23K 1/17, A23L 3/3571, A61K 37/54, C12Q 1/04

The microorganism(s) has (have) been deposited with NCTC under numbers 12452, 12453 and with NCIMB under number 40400.

- 30 Priority: 20.04.91 GB 9108498
- (43) Date of publication of application: 28.10.92 Bulletin 92/44
- (A) Designated Contracting States:

 AT BE CH DE DK ES FR GB GR IT LI LU MC NL
 PT SE
- (1) Applicant: AGRICULTURAL AND FOOD RESEARCH COUNCIL Norwich Research Park Colney, Norwich NR4 7UA (GB)
- (2) Inventor: Gasson, Michael John Old Ash Farm, Stone Lane, Bintree Dereham, Norfolk NR20 5NA (GB)
- (4) Representative: Bassett, Richard Simon et al ERIC POTTER & CLARKSON St. Mary's Court St. Mary's Gate Nottingham NG1 1LE (GB)
- (A) Bacteriophage lysins and their applications in destroying and testing for bacteria.
- Bacteriophages of food-contaminating or pathogenic bacteria or the lysins thereof are used to kill such bacteria. Examples include lysins from bacteriophages of *Listeria monocytogenes* and *Clostridium tyrobutyricum*.

Tests for bacterial contamination can be made specific for specific bacteria by using the appropriate bacteriophage or lysin thereof and determining whether cells are lysed thereby.

This inventin relates to the use of bacterial viruses (bacteriophages) which use bact ria as hosts and produce a bact riophage lysin responsible for cell-wall degradation and lysis of the host cells.

Attempts to use a bacteriophage as an antimicrobial agent have failed to be effectiv. We have previously used the lysin of the bacteriophage øvML3 of Lactococcus lactis ML3, which is active against all strains of all subspecies of Lactococcus lactis, very weakly affects group Denterococci, but does not have any action on a wide variety of other species tested (Shearman et al (1989) Molecular and General Genetics 218: 214-221), to lyse cheese starter cultures (WO90/00599). WO/00599 also discloses the use of micro-organisms, transformed to express the øvML3 lysin, to suppress populations of bacteria susceptible to the lysin, ie the Lactococcus lactis cheese starter culture strains.

It is also known to use cheese starter culture bacteria to produce the simple peptide nisin in order to destroy harmful bacteria.

We have now found that further bacteriophage lysins can be used to destroy unwanted bacteria, especially food-contaminating bacteria prejudicial to health.

A first aspect of the present invention provides a formulation comprising a lysin of a bacteriophage of a food-contaminating or pathogenic bacterium or a variant of such a lysin, substantially free of the bacteriophage.

Preferably the *Listeria* phage øLM4 or *Clostridium tyrobutyricum* phage øP lysins are used. They act against all tested species and strains of *Listeria* and also strains of *Kurthia zopfii*, or against *Clostridium tyrobutyricum* (as appropriate), but lack activity against other tested species.

A "variant" of such a lysin is any polypeptide of which at least 30% (preferably at least 50%, 75%, 90%, 95% or 99%) has at least 80% (preferably at least 90%, 95% or 99%) amino acid homology with the corresponding region of the lysin itself and which has at least 30% (preferably at least 50%, 75%, 90% or 95%) of the bacterial lysing capability of the said lysin.

Food-contaminating bacteria are those which, by virtue of their presence or compounds produced by them, cause undesirable flavours, odours or visual appearances or cause illness in humans or animals consuming the food.

The organism which is destroyed may be any of the following:
Listeria monocytogenes, Clostridium tyrobutyricum, Clostridium botulinum, Clostridum perfringens, lactic acid bacteria (eg Lactobacillus brevis) causing beer spoilage, Salmonella spp., Yersinia spp., Campylobacter, E. coli, Pseudomonas spp., Staphylococcus, Bacillus spp. (including Bacillus cereus), Shigella spp. and Vibrio spp.

Pathogenic bacteria include all pathogenic bacteria of humans, animals and plants. However, in a medical or veterinary context, as is explained further below, bacteria involved in topical or superficial infections are of particular interest. These include Staphylococcus spp. (eg Staph. aureus), Streptococcus spp., Corynebacterium spp., Clostridium spp. (eg Cl. perfringens), Yersinia spp. (eg Y. pestis), Pasteurella spp. (eg P. multocida), Streptobacillus spp. (eg Streptobacillus moniliformis), Proteus spp. (eg P. mirabilis) and Pseudonomas spp.

A second aspect of the invention provides a substantially pure preparation of a lysin from a bacteriophage of a food-contaminating or pathogenic bacterium.

A third aspect of the invention provides a coding sequence for such a lysin.

A fourth aspect provides a DNA construct comprising a coding sequence as above in an expression vehicle suitable for transformation of a microbial host or cell line.

Suitable regulatory expression vectors, transformation techniques, and hosts are all known in the art. The host may be any micro-organism or cell line which is found to express the said lysin gene, and may be a bacterium such as *E. coli* or *Lactococcus lactis*, a yeast such as *Saccharomyces cerevisiae* or *Kluveromyces lactis* or a filamentous fungus such as *Asperaillus niger*.

Thus a fifth aspect provides a microbial or cell line host transformed with such an expression vehicle and capable of expressing the lysin coding sequence.

A sixth aspect provides a polypeptide derived from the expression of the said lysin coding sequence in a suitable host transformed with such an expression vehicle.

A seventh aspect of the present invention provides a method of destroying pathogenic or food-contaminating bacteria characterised in that said bacteria are lysed with a lysin or a variant of such a lysin from a bacteriophage of such bacteria.

The use of such a preparation in food or agriculture simply involves the addition of an amount sufficient to provide an inhibitory concentration of lysin activity. The specific activity of any preparation may readily be calculated, for example by use of the spectrophotometric assay described later. The quantity of preparation necssary for effective protection in a given f od may be arrived at by reutine experimentation. The lysin is applied in a suitable, non-toxic aqueous medium. Any food may be treated with such a preparation by addition or application to surfaces eg of cut, cooked meat or poultry, soft cheeses and pâtés of fisher meat. The term "food" includes drinks (such as water, ber, milk and soft drinks), animal food (such as pet food or cattle food) and

10

20

25

30

produce destined fir consumption by humans or animals (such as stored potatoes). In agriculture, a particular application is addition to silage wher Listeria and Clostridium tyrobutyricum are known to present a problem that can be passed on up the food chain. In briwing, brewing yeast transfirmed with a lysin gene may be used.

In a medical or veterinary context, b caus the lysin is lik ly to be d graded or to produce an immune reaction, it is preferred to administer it topically in diseases of the skin such as ulcers, burns and acne. It may be applied as the clinician directs, as a lotion, cream or ointment.

An eighth aspect provides a method of testing for the presence of bacteria which are lysed by a bacteriophage or by the lysin thereof, comprising exposing a sample to the said bacteriophage or lysin and determining whether bacteria have been lysed as a result of such exposure.

Any technology that exploits the release of intracellular biochemicals (eg ATP or enzymes such as alkaline phosphatase or esterase) to detect micro-organisms can, in accordance with the invention, be made specific for the target range of such lysins. For example, an ATP or phosphatase release test for *Listeria* using the *Listeria* bacteriophage or lysin thereof, in which the release of ATP or phosphatase is detected (eg by linkage to a luciferase reaction and monitoring of photon release or by spectrophotometric methods as is described below) indicates the specific presence of *Listeria* in a sample. The invention further provides a kit comprising a lysin and means to detect bacterial lysis.

Preferably, the bacteriophage in all these contexts is or at least includes *Listeria monocytogenes* øLM4 or a bacteriophage of *Clostridium tyrobutyricum*, such as øP1. Several different lysins may be used in order to destroy or identify a specific range of bacteria.

The cloning and characterization of the gene for the lysin of the *Listeria* bacteriophage øLM4 has facilitated the production of the free lysin and the availability of its structural gene. These components have application in the protection of environment and food material from pathogenic strains of *Listeria*. The free lysin acts as a novel antimicrobial that kills such bacteria and the gene can be genetically engineered in a non-pathogenic micro-organism such that the latter produce the *Listeria* lysin thereby equipping it with a novel anti-*Listeria* capability. For example, a food-grade micro-organism may be transformed with a DNA construct comprising a coding sequence for the lysin.

Preferred embodiments of the invention will now be described by way of example with reference to the accompanying drawings, in which:-

Figure 1 shows patches of *E. coli* clones with *Hin*dIII fragments of øLM4 DNA in the *Hin*dIII site of vector pUC18. The plate is overlayed with a suspension of *Listeria monocytogenes* 6868 cells and lysin producing clones create clear zones around the patch (indicated by an arrow).

Figure 2 is a restriction and deletion map of lysin-expressing done pFl322. The result of lysin activity tests is indicated to the right. The inferred location of the lysin gene is shown. Arrows indicate the orientation of the lysin gene with respect to the lac α promoter of the pUC vector used which is transcribed from left to right in this figure (ie pFl324 is opposed to the lac α promoter, other clones are transcribed in the same direction as the lac α promoter).

Plasmid pF1322 is pUC18 carrying a 3.6kb HindIII fragment of bacteriophage øLM4 DNA. Plasmid pF1326 is pF1322 with a 0.56kb HindIII - Sall deletion. Plasmid pF1327 is pF1322 with a 1.32kb HindIII - EcoRI deletion. Plasmid pF1324 is pUC18 carrying a 1.9kb HindIII - Nrul fragment of pF1322 cloned between its HindIII and HincII sites. Plasmid pF1325 is pUC18 carrying a 1.6kb Nrul - HindIII fragment of pF1322 cloned between its HindIII sites. Plasmid pF1328 is pUC19 carrying a 1.9kb HindIII - Nrul fragment of pF1322 cloned between its HindIII and HincII sites. Plasmid pF1329 is pF1328 carrying a 1.6kb Baßl deletion from the polylinker BamHI site. Plasmid pF1330 is pF1328 carrying a 1.6kb Baßl deletion from the polylinker BamHI site.

Figure 3 illustrates the response of a suspension of *Listeria monocytogenes* 6868 cells to cell free extracts of *E. coli* strains harbouring plasmids pFl322(♠), pFl328(△), pFl329(○) and pUC19(●).

Figure 4 is a Coomassie blue stained SDS polyacrylamide gel of proteins produced by *E. coli* strain carrying the T7 expression vector pSP73 (tracks 2 and 3) or pFl331 which carries the lysin gene (tracks 4 and 5). Uninduced cells (tracks 2 and 4) are compared with induced cells (tracks 3 and 5). Molecular weight markers are present (tracks 1 and 6) and the expressed lysin protein is indicated by an arrow.

Figure 5 illustrates the sequencing strategy used. The extent and direction of sequences determined are indicated by the arrows. Synthetic oligonucleotide primers are indicated by boxes.

Figure 6 shows a single strand of the region of øLM4 DNA that encodes the lysin gene.

Figure 7 is the Analyseq print out of the analysis of the DNA sequence shown in Figure 6. The identification of the open reading frame of the lysin gene is in the top panel.

Figure 8 sh ws the double stranded DNA sequence of the lysin structural gene and its translat id protein product.

Figur 9 shows the prot ctive effect f clon d Listeria lysin on skimmed milk to which Listeria Monocytogen s is added.

45

50

55

15

Figure 10 shows the expression of the *Listeria* lysin gene in *Lactococcus lactis* under the control of the lactose inducible lact coccal lactose operon promoter.

EXAMPLE 1: CLONING OF LYSIN GENE, ETC

Isolation of bacteriophage øLM4

5

20

30

A bacteriophage named øLM4 was isolated from a culture of *Listeria monocytogenes* serotype 4b that was originally obtained from a listeriosis outbreak in Nova Scotia, Canada in 1981. The source of the infection was tracked down to contaminated coleslaw. This culture of *Listeria monocytogenes* was deposited under the Budapest Treaty as NCTC 12452 in the National Collection of Type Cultures, Central Public Health Laboratory, Colindale, London, UK on 21 March 1991. The bacteriophage was purified by standard single plaque isolation procedure using *Listeria monocytogenes* F6868 as the host. This culture was similarly deposited under the Budapest Treaty as NCTC 12453 in the National Collection of Type Cultures, Central Public Health Laboratory, Colindale, London, UK on 21 March 1991. Examination of this bacteriophage by electron microscopy revealed it to have an isometric head with a diameter of approximately 50nm and a tail of approximately 250nm.

Isolation of DNA from Listeria monocytogenes bacteriophage øLM4

20ml of an 18 hour culture of *Listeria monocytogenes* F6868 was inoculated into 500ml of Bacto tryptose phosphate broth and incubated with shaking at 30°Co When O.D. 600 reached 0.15 the culture was infected with 5 x 10¹⁰ p.f.u. of bacteriophage eLM4 and incubated until lysis was apparent as a loss of turbidity. The lysate was centrifuged at 6000 x g for 10 min at 4°C. The bacteriophage lysate was then concentrated by polyethylene glycol precipitation and purified on caesium chloride stepped gradients using well established protocols (Bachrach and Friedmann (1971) *Applied Microbiology* 22: 706-715). Bacteriophage DNA was extracted by dialysis against 50% formamide in TE buffer (0.1 Tris - HCI, 0.01M EDTA, pH 8.5). Further purification was then performed on caesium chloride-ethidium bromide equilibrium density gradients. Examination of the bacteriophage DNA by agarose gel electrophoresis revealed the genome to be approximately 39kb in size.

Cloning the bacteriophage sLM4 lysin gene

DNA purified from bacteriophage eLM4 was digested with restriction endonuclease *Hin*dIII and ligated to plasmid pUC18 vector DNA that had also been cleaved with restriction endonuclease *Hin*dIII. The ligated DNA was transformed into *Escherichia coli* TB1 and ampicillin resistant colonies were selected on LB agar containing 50μg/ml ampicillin, 40μg/ml isopropyl-β-D-thiogalactopyranoside (IPTG) and 40μg/ml 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-gal). These steps were performed using well established protocols (Sambrook, J. *et al* (1989), *Molecular Cloning. A laboratory manual.* Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 2nd Edition).

White colonies were screened for their ability to produce a bacteriophage lysin active against *Listeria monocytogenes*. These colonies were patched onto duplicate Bacto tryptose agar plates and incubated for 18 hours at 37°C. One plate was exposed to chloroform vapour for 10 min and then seeded with 0.2ml of an 18 hour broth culture of *Listeria monocytogenes* F6868. After incubation at 30°C for 18 hours clear zones of lysins were apparent around patches of clones expressing the *Listeria* bacteriophage øLM4 lysin. This is illustrated in Figure 1. Positive clones were recovered from the duplicate plate and the pUC18 derivative plasmid isolated and characterized by digestion with restriction endonuclease *Hind*III. One lysin expressing pUC18 clone that contained a 3.6kb insert of øLM4 DNA was chosen for further analysis. This plasmid was designated pF1322.

Deletion analysis of lysin expressing plasmid pF1322

Characterization of pFI322 was undertaken by constructing a restriction map of this insert using single and double digests with a variety of restriction enzymes. The map is presented in Figure 2. Deletion of some regions of the 3.6kb insert contained in pFI322 was achieved by digestion with certain of these enzymes, religation and transformation into *E. coli* TB1. In other instances endonuclease *Bal 31* was used to introduce deletions. In addition, some regions of the 3.6kb cloned DNA in pFI322 were deleted by digestion with certain restriction endonucleases and re-cloning into appropriately cleaved plasmid vectors pUC18 or pUC19 and transformation into *E. coli* TB1. These manipulations are clearly documented in Figure 2 which is presonted in the form of a document of the product of the

as the plate assay described above and illustrated in Figure 1, a spectrophotometric assay was also us d. For this th *E. coli* strain carrying plasmid clones wer grown at 37°C f r 18 hours, harvested by c ntrifugation at 6000 x g f r 5 min at 4°C, washed down nce in 100mM Tris buff r pH7.5 and resusp nded in this same buffer at approximately 10mg dry weight/ml. Cell free extracts were mad by 6 cycles of ultrasonication (15 sec on, 10 sec off) at 0°C using the microprobe of an MSE Soniprep 150. Unbroken cells and cell debris were removed by centrifugation at 25000 x g for 15 min at 4°C.

Samples of the cell free extracts were added to an equilibrated (5 min at 37°C) 4ml reaction mixture containing 400µmole Tris HCl pH7.5 and *Listeria monocytogenes* F6868 indicator cells that had been harvested and resuspended at an O.D. 600 of 2.3. The fall in optical density caused by lysis of indicator cells was followed using a spectrophotometer. Typical results from use of this protocol are presented in Figure 3. The lytic activity of the plasmid derivative described above and in Figure 2 were assessed using both of these methods and the results are presented in Figure 2.

These results demonstrated that the structural gene for bacteriophage øLM4 was contained within the left hand 1.2kb of the DNA cloned in pFl322 and defined by the *HindIII* site at co-ordinate 0 and the *EcoR1* site at co-ordinate 1.25 of the map illustrated in Figure 2.

Figure 2 also indicates the orientation of Listeria bacteriophage \varnothing LM4 DNA with respect to the E. coli lac α promoter that is present on vectors pUC18 and pUC19. It is apparent that a positive reaction in the lysin assay is only found when one orientation is maintained (eg pFl324 is negative whereas pFl328 is positive even though both constructs contain the same Listeria bacteriophage \varnothing LM4 fragment). This suggests that expression of the lysin gene depends on use of the E. coli lac α promoter and that no Listeria bacteriophage \varnothing LM4 promoter is present and active in E. coli.

Detection of the lysin protein

In order to identify a protein produced by the fragment of øLM4 DNA that expressed lysin activity another *E. coli* vector was used. A 2kb fragment from plasmid pFl328 between the *HindIII* site at co-ordinate 0 and a unique *BamHI* site present on the polylinker of pUC19 was isolated and cloned between the *HindIII* and *BamHI* sites of the T7 expression vector pSP73 that was purchased from Promega. The constructed plasmid named pFl331 was transformed into the *E. coli* host strain JM109DE3.

The *E. coli* T7 promoter in this vector is expressed by the phage specific T7 RNA polymerase which is induced by addition of IPTG in the appropriate host strain *E. coli* JM109 DE3. Cultures of this strain carrying pSP73 as a control or pFl331 were grown for 3 hours and induced by addition of 1PTG to a final concentration of 0.2mM. Incubation was continued for a further 3 hours before the cultures were harvested and used to prepare cell extracts using well-established, published procedures (Studier, F.W., Rosenberg, A.H., Dunn, J.J. and Dubendorff, J.W. (1990) *Methods in Enzymology* 185: 60-89).

Proteins present in cell extracts were analysed using conventional SDS-polyacrylamide gel electrophoresis (Laemmli (1970) *Nature* **227**: 680-685). The results presented in Figure 4 clearly demonstrate that the 2kb fragment of pFi331 expresses a single protein with a molecular size of 31 kilodaltons which represents the lysin enzyme.

-

DNA sequence of the Listeria bacteriophage øLM4 lysin gene

The region of DNA between co-ordinate 0 and 1.2 in Figure 2 was subject to oligonucleotide sequence analysis using the dideoxy chain-termination method (Sanger, F., Coulson, A.R., Barrell, B.G., Smith, A.J.H. and Roe, B.A. (1980) *J. Molec. Biol.* 143) with a sequenase version 2.0 kit (United States Biochemical Corporation). The 0.9kb *HindIII - EcoRI* and the 0.3kb *EcoRI - EcoRI* fragments of pF1328 were subcloned in the M13 sequencing vectors M13mp18 and M13mp19 to create templates and sequenced using universal and synthetic oligonucleotide primers. To sequence across the internal *EcoRI* site at co-ordinate 0.9 double stranded sequencing of pF1329 plasmid DNA was used. The sequencing strategy is presented in Figure 5 and the complete DNA sequence is in Figure 6. The sequence was analysed using the computer programme ANALYSEQ (Staden (1980) *Nucleic Acid Research* 8: 3673–3694) which revealed an open reading frame that represents the *Listeria* bacteriophage lysin gene. The printout from the Analyseq analysis is presented in Figure 7 and the open reading frame representing the lysin structural gene and its translated protein product is presented in Figure 8. The molecular size of the translated protein was calculated to be 32.9 kilodaltons which agrees well with the calculated 31 kilodalton size of the protein express d by the T7 vector pSP73 (Clone pF1331 in Figure 4).

Activity and spe ificity f the List ria bacteriophage øLM4 lysin

Figure 3 illustrates the lytic activity of crude cell free extracts of *E. coli* TB1 carrying the plasmids pFl322, pFl328, pFl329 and pUC19 assayed using the spectroph to metric method described above. This activity was related to units of commercially available mutanolysin (Sigma) as has been described previously (Shearman, C., *Underwood*, H., Jury, K. and Gasson M. (1989) *Mol. Gen. Genetics* 218: 214-221). The crude cell extracts of lysin expressing clones typically contained 5000 mutanolysin equivalent units per mg. protein.

In order to test the spectrum of activity of this lysin, the spectrophotometric assay was performed on 16 serotypes of *Listeria monocytogenes*, all other species of *Listeria*, the related species *Kurthia zopfii* and a variety of other gram positive and gram negative bacteria. The results compiled in Table 1 show that the *Listeria* bacteriophage eLM4 lysin was active against all tested strains of *Listeria monocytogenes*, *Listeria innocua*, *Listeria ivanovii*, *Listeria murrayi*, *Listeria seelegri*, *Listeria welshimeri*, *Listeria grayi* and *Kurthia zopfii*. No activity was found against any of the other species tested.

TABLE 1: ACTIVITY OF CLONED LYSIN AGAINST LISTERIA SPECIES

5	Organism	Strain	Serotype	Relative!	Time (min)b
				Activity	ΔOD ₆₀₀ =1
		.		-	,
10	Listeria	F6868	4b	1.00	20
	monocytogenes	NCTC 7973	1a	0.19	53
		NCTC 5412	4b	0.90	13
		F4642	4 b	0.92	14
15		NCTC10357	1a	0.92	20
		BL87/41	4	0.66	25
		NCTC 5348	2	0.10	78
20		SLCC2373	3a.	1.20	17
		SLCC2540	3 b	0.19	60
		SLCC2479	3c	0.15	60
25		SLCC2374	4a	0.54	30
		SLCC2376	4c	0.19	90
		SLCC2377	4 d	0.08	90
		SLCC2378	4e	0.56	28
30		SLCC2482	7	0.45	36
		L3056	1/2a	0.49	30
		L4203	1/2a	0.36	41
35					
	Organism	Strain	Serotype	Relative*	Time (min)b
				Activity	ΔOD ₆₀₀ =1
40					
		L4490	1/2b	0.29	55
		L1378	1/2b	0.09	150
		L4281	1/2c	0.11	120
45		L3304	1/2c	0.12	90
		L3253	4bx	0.66	26
		L2248	4bx	0.08	72
50	Listeria	NCTC11288	6a	0.90	12
	innocua				
		NCTC11289	6a	0.69	22
55	Listeria	NCTC11007		0.95	18
55	ivanovii				

		SLCC5579	0.51	30
	Listeria	NCTC11856	1.10	15
5	seeligeri			
	Listeria	NCTC11857	0.29	36
	welshimeri			
	Listeria	NCTC10812	0.86	15
10	murrayi			
	Listeria	NCTC10815	0.93	12
	grayi			
15	Kurthia	NCTC10597	0.54	28
	zopfii			

Table 1 shows the relative sensitivity of a selection of strains of *Listeria* and *Kurthia zopfii* to the bacterio-phage øLM4 lysin. a) Relative activity is the fall in optical density (O.D₆₀₀) from 2.3 achieved in 30 minutes divided by the equivalent fall obtained using *Listeria monocytogenes* F6868. b) The time (min) taken for a fall in optical density of O.D.₆₀₀ from 2.3 to 1.3 (O.D.₆₀₀ fall of 1) is recorded. Other strains tested which show no sensitivity to lysin were *Aeromonas hydrophila* NCTC 8049, *Bacillus cereus* NCTC 11143, *Brocothrix thermosphacta* NCTC 10822, *Camobacterium pisciola* BL90/14, *Enterococcus faecalis* BL90/11, *Escherichia coli* BL90/12, *Klebsiella pneumoniae* NCFB 711, *Pseudomonas fluorescens* BL 78/45, *Staphylococcus aureus* NCTC 10652, *Streptococcus pneumoniae* NCTC 7465, *Streptococcus pyogenes* NCTC 2381.

In addition it was observed that the lysin was active at temperatures as low as 2°C. At 2°C addition of lysin to suspensions of *Listeria monocytogenes* caused a decrease of between 0.7 and 2.0 0.D.₆₀₀ units within 24 hours.

EXAMPLE 2: USE OF LYSIN TO CONTROL LISTERIA

Use as a free lysin

30

35

There are two distinct application concepts. One exploits a preparation of lysin enzyme manufactured by fermentation of a genetically engineered micro-organism that expresses the lysin gene product (Free lysin). The host organism may be *E. coli*, or any other bacterial species such as *Lactococcus lactis*, a yeast such as *Saccharomyces cerevisiae or Kluveromyces* lactis or a filamentous fungus such as *Aspergillus niger*. The lysin gene may be expressed intracellularly in which case a preparation may consist of a cell free lysate of the producing organism with some purification of the lysin, for example by ammonium sulphate precipitation and/or column chromatography. Alternatively the fermentation micro-organism may secrete lysin into the culture medium in which case the supernatant of the centrifuged fermentation broth provides the basis of a preparation, which again may require some purification.

The effectiveness of a crude extract of cloned Listeria lysin was demonstrated by its addition to skimmed milk containing *Listeria monocytogenes*. As illustrated in Fig. 9 the lysin preparation reduces the viable count of *Listeria monocytogenes* and after 22 days incubation at 8°C there is a viable count difference of 10⁸ *Listeria* cfu between milk containing lysin and the control sample.

Expression of Lysin by a genetically engineered micro-organism

An alternative application concept is to use a genetically engineered micro-organism that is compatible with a food or agricultural environment such as a species of lactic acid bacteria. Such an organism then grows in a food or agricultural environment and expresses an introduced gene for *Listeria* bacteriophage lysin. The gene may be xpr ssed intracellularly and releas d into food or an agricultural nvir nment by autolysis or induced lysis of that micro-organism. Alt matively the lysin may be secreted by a micro-organism so that active lysin is reliased dinto a food in agricultural environment by that viable micro-organism. In these cases the lysin gene is placed downstream of an appropriate promoter such as the lactose operon promoter or the proteinas promot in of *Lactoc ccus lactis* NCFB 712. So critical may be achieved by fusion of the lysin structural gene to a

known N t minals cretory leader such as those of the proteinase gene, the *usp*45 g ne r the nisin precursor gen of *Lactococcus lactis*. Suitable organisms for this application concept include strains of *Lactococcus lactis* in che se and dairy products and *Lactobacillus plantarum* or *Pediococcus* sp cies in agricultural silage.

The Listeria lysin gen from plasmid pF1328 was isolated together with its wn ribos me binding site using the polymerase chain reaction. This fragment was cloned into the PstI site of *E. coli* vector pUC19 in both orientations (plasmids pF1531 and pF1532). Expression of this gene in *E. coli* strains was observed from one orientation only, under the control of the *lac* α promoter of the vector (plasmid pF1531). Enzyme activity of cell extracts of this strain was comparable to that of *E. coli* strains carrying plasmid pF1531. Enzyme activity of cell extracts of this strain was comparable to that of *E. coli* strains carrying plasmid pF1532. Using plasmid pF1532 that did not express the lysin gene and cloning the lactococcal *lac*A promoter/lacR gene on a *Bam*HI fragment (Van Rooijen *et al.* (1992) *J. Bacteriol.* 174: 2273-2280) upstream of the lysin gene (plasmid pF1533) expression in *E. coli* of eLM-4 lysin from the lactococcal lacA promoter was obtained. The lytic activity of extracts from these *E. coli* strains was lower when the lysin gene was expressed from the lacA promoter. The *Ssti/SphI* fragment of pF1533 containing the eLM-4 lysin gene with the lacA promoter/lacR gene was cloned into the *Sstl/SphI* sites of the lactococcal vector pTG262 (Shearman *et al* (1989) *Molecular and General Genetics* 218: 214-221) and the resulting plasmid pF1534 was used to transform *L. lactis* MG5267. As shown in Figure 10 cell extracts of this strain expressed eLM-4 lysin activity when grown on lactose, on glucose enzyme activity of cell extracts was reduced.

The øLM-4 lysin gene together with the lacA promoter/lacR gene was cloned into pF145, a plasmid expressing the *Lactococcus* phage øvML3 lysin gene which causes lysis during stationary phase of *L. lactis* cultures carrying the plasmid (Shearman et al (1992) *Biotechnology* 10: 196-199). The resulting plasmid pF1535 in *L. lactis* MG5267 when grown on lactose produced a culture that grew to stationary phase, then lysed as a consequence of the øvML3 lysin, releasing øLM-4 lysin into the culture supernatant.

EXAMPLE 3: SPECIFIC DETECTION OF MICRO-ORGANISMS

25

The specificity of a bacteriophage lysin provides an opportunity to specifically detect those micro-organisms which are susceptible to it. For example to detect *Listeria sp.* the lysin described here may conveniently be used at a post enrichment stage where a broth culture of those micro-organisms present in a test sample is first produced. The identity of species of bacteria in the sample at this stage is unknown. The bacterial culture may be centrifuged and resuspended in an assay buffer (eg the one used here in studies of lysin specificity). A control preparation and separately a preparation containing active *Listeria* lysin are then added. Sufficient units of lysin activity are used to provide very effective lysis of any lysin susceptible cells (ie *Listeria*). After incubation for a short period (eg 30 min) any *Listeria* present will lyse, but other species will not. The presence of *Listeria* will then be detected by the lysis of bacteria in the sample treated with the lysin whereas no lysis occurs in the control.

The detection of lysis may be achieved by assaying an intracellular enzyme or metabolite. Especially useful enzyme assays are for phosphatase or for esterase. Alkaline phosphate can be assayed spectrophotometrically by following appearance of p-nitrophenol, which is yellow, from the colourless substrate p-nitrophenyl-phosphate at 405nm. Esterase activity can be assayed using fluorescein diacetate which is cleaved to acetate and fluorescent fluorescein and measuring the latter in a fluorometer. One especially suitable metabolite assay involves ATP detection. For this the well established luciferase assay in which ATP molecules generate light is exploited. Light emission may be measured in a luminometer. (An example of an end point detection reagent using luciferase-luciferin is marketed by Sigma Chemical Company as product L-1761).

EXAMPLE 4: CLOSTRIDUM TYROBUTYRICUM BACTERIOPHAGE ®P1 LYSIN

Bacteriophage @P1 was isolated from a landfill core sample using Clostridium tyrobutyricum NCFB 1755 as host. Bacteriophage @P1 was tested against six more strains of C. tyrobutyricum. Strains NCFB 1753 and NCFB 1756 supported the growth of bacteriophage and they were thus host strains as was the strain NCFB 1755. Against C. tyrobutyricum strains NCFB 1715, NCFB 1754, NCFB 1757 and NCFB 1790 an undiluted @P1 stock suspension gave a clear zone but diluting out did not result in individual bacteriophage plaques. This indicates that these strains were lysin sensitive but not bacteriophage sensitive. Bacteriophage @P1 thus produces a lysin with a broad specificity for strains of C. tyrobutyricum. Similar tests of bacteriophage @P1 with a wide variety of other bacteria showed no effect of the lysin or bacteriophage particles against C. sporogenes strains ATCC 17886, NCFB 1789, NCFB 1791; C. butyricum strains NCFB 1713, NCFB 857; Lactobacillus buitinen strains NCFB 110, F3327; L. brevis strains NCFB 1749, F3328; L. helveticus strains NCFB 1243, CNRZ 832; L. bulgaricus CNRZ448; L. plantarum strains NCFB 1752, NCFB 82, NCFB 963; Escherichia coli BL 90/12; Bacillus cereus NCTC 1143.

Bact riophage eP1 was deposited at the National Collections of Industrial and Marine Bacteria, 23 St Machar Drive, Ab rde n, AB2 1RY, Scotland on 5 April 1991 and a new deposit was made on 4 July 1991 under the Budap st Treaty and has been accorded Accession No NCIMB 40400.

Claims

5

10

- A formulation comprising a lysin of a bacteriophage of a food-contaminating or pathogenic bacterium or a variant of such a lysin, substantially free of the bacteriophage itself.
- 2. A formulation according to Claim 1 wherein the bacteriophage is a Listeria monocytogenes bacteriophage.
- 3. A formulation according to Claim 2 wherein the bacteriophage is øLM4.
- 4. A formulation according to Claim 1 wherein the bacteriophage is a Clostridium tyrobutyricum bacteriophage.
 - A method of destroying micro-organisms characterised in that said micro-organisms are lysed with a formulation according to any one of the preceding claims.
- A method according to Claim 5 wherein pathogenic strains of Listeria or Clostridium are destroyed in or on food products.
 - 7. A substantially pure preparation of a Listeria or Clostridium bacteriophage lysin.
- 8. A nucleotide coding sequence for the lysin of a bacteriophage of a food-contaminating or pathogenic bacterium or a variant of such a lysin.
 - A coding sequence comprising the DNA coding sequence given in Figure 6 or a variant thereof encoding the same polypeptide.
- 10. An expression vehicle comprising a coding sequence according to Claim 8 or 9 and regulatory regions associated therewith for expression of the coding sequence in a suitable host.
 - 11. A microbial host transformed with means to express a lysin of a bacteriophage of a pathogenic or food-contaminating bacterium or a variant of such a lysin.
 - 12. A host according to Claim 11 which is a food-grade micro-organism.
 - 13. A lysin derived from the cultivation of a host according to Claim 11:
- 40 14. A method of testing for specific bacteria in a sample, comprising adding a bacteriophage lysin to the sample and determining whether bacterial cells have been lysed thereby.

50

45

35

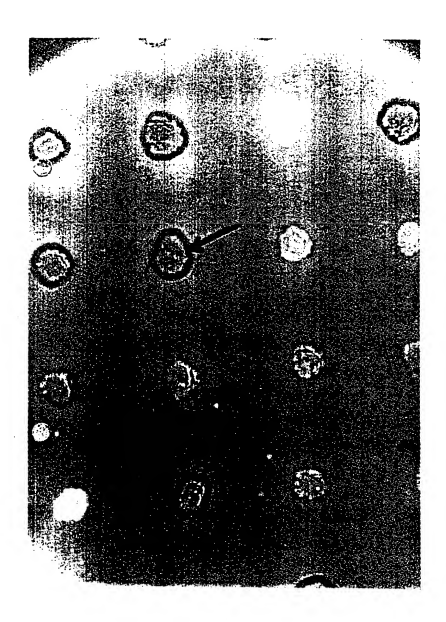


Fig. 1

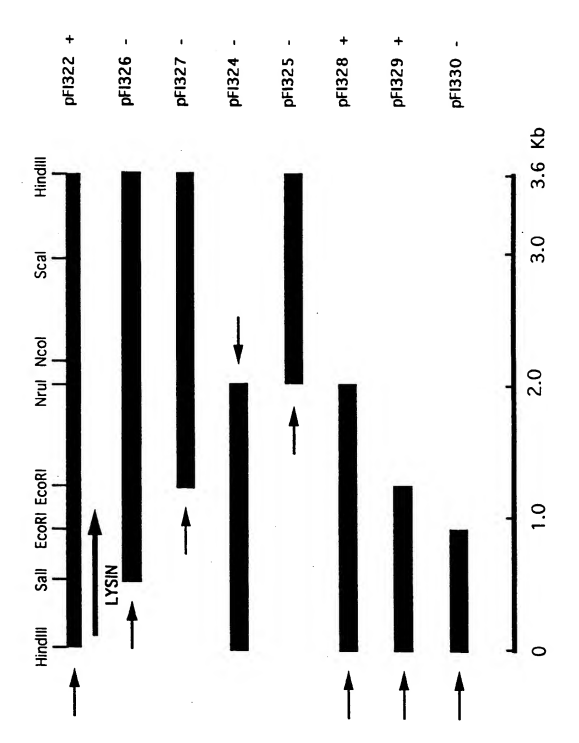


FIGURE 2

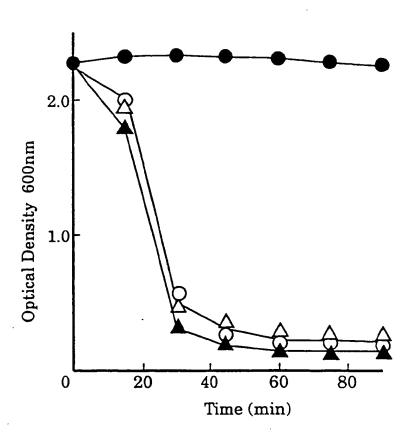


FIGURE 3

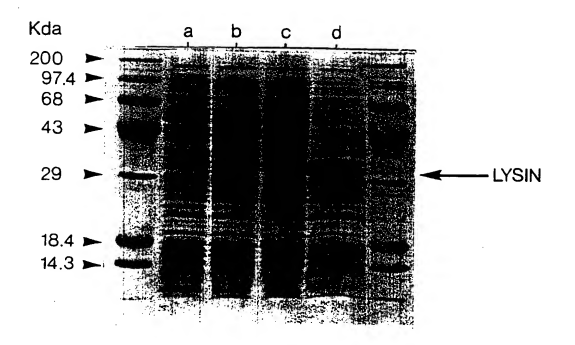


Fig. 4

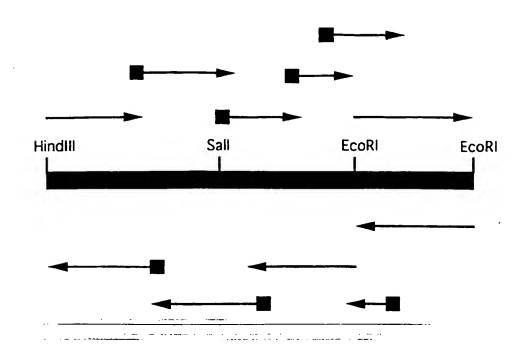


FIGURE 5

1	AAGCTTTACT	GGATAACGAC	AAACTAGĀÁC	TGACAGATGA	AGGACTGAAA
51	GGCCTTGACG	AACAGCTAGG	AGCATTGCAA	GAAAGCGATG	CTTATTTGTT
101	TGCTCAAGAA	AGCGAGGCGG	GGCGGAACTG	GTTTATTTGA	ACAATTTACT
151	AATCGAGCTA	AAAAATATGG	AAAGGATGAT	TAATAATGGC	ATTAACAGAG
201	GCATGGCTAA	TTGAAAAAGC	AAATCGCAAA	TTGAATACGT	CAGGTATGAA
251	TAAAGCTACA	TCTGATAAGA	CTCGGAATGT	AATTAAAAA	ATGGCAAAAG
301	AAGGGATTTA	TCTTTGTGTT	GCGCAAGGTT	ACCGCTCAAC	AGCGGAACAA
351	AATGCGCTAT	ATGCACAAGG	GAGAACCAAA	CCTGGAGCGA	TTGTTACTAA
401	TGCTAAAGGT	GGGCAATCTA	ATCATAATTT	CGGTGTAGCA	GTTGATTTGT
451	GCTTGTATAC	GAGCGACGGA	AAAGATGTTA	TTTGGGAGTC	GACAACTTCC
501	CGGTGGAAAA	AGGTTGTTGC	TGCTATGAAA	GCGGAAGGAT	TCGAATGGGG
551	CGGAGATTGG	AAAAGTTTTA	AAGACTATCO	GCATTTTGAA	CTATGTGACG
601	CTGTAAGTGG	TGAGAAAATC	CCTACTGCGA	CACAAAACAC	CAATCCAAAC
651	AGACATGATG	GGAAAATCGT	TGACAGCGC	CCACTATTGC	CAAAAATGGA
701	CTTTAAATCA	AATCCAGCGC	GCATGTATA	ATCAGGAACT	GAGTTCTTAG
751	TATATGAACA	TAATCAATAT	TGGTACAAGA	A CGTACATCA	CGACAAATTA
801	TACTACATGT	ATAAGAGCTT	TTGCGATGT	T GTAGCTAAA	AAGATGCAAA
851	AGGACGCATC	AAAGTTCGAA	TTAAAAGCG	G GAAAGACTT/	CGAATTCCAG
901	TTTGGAATAA	CACAAAATT	AATTCTGGG	A AAATTAAAT	G GTATGCACCC
951	AATACAAAAT	TAGCATGGT	A CAACAACGG	A AAAGGATAC	T TGGAACTCTG
1001	GTATGAAAAG	GATGGCTGG	T ACTACACAG	C GAACTACTT	C TTAAAATAAA
1051	AAGTCCCGGT	TTGAGCTGG	G CTTTTTATT	T TGAAAGTGA	C TAACAAAAA
1101	TGTAATAAA	A ATGTAATAA	T CCAAGTAAG	T TGTATAAAA	T TTGCAGAATT
1151	AGAACGTTTT	T ATTGATAAA	T ACACCTTAT	G AAGAGTATT	T TTGGCTATAT
1201	TTGCGCATT	A TAGGGTTGA	A TGTAACACT	A TATGTAGAA	T TC

FIGURE 6

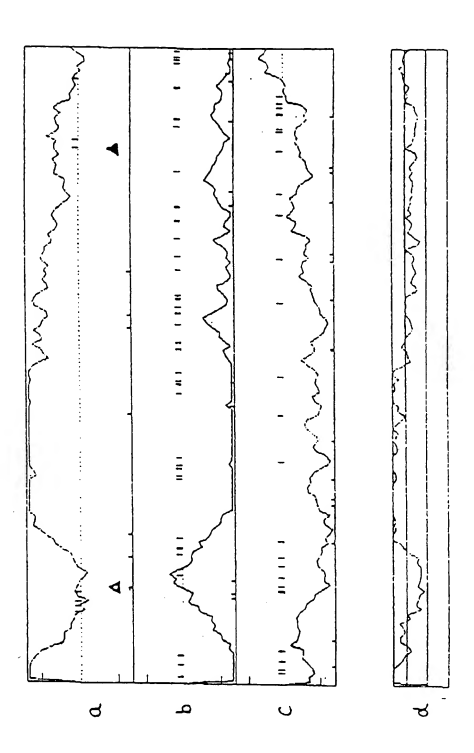


FIGURE 7

106	ATGGCATTAACAGAGGCATGGCTAATTGAAAAAGCAAATCGCAAATTGAATACGTCAGGT	245
186	TACCGTAATTGTCTCCGTACCGATTAACTTTTTCGTTTAGCGTTTAACTTATGCAGTCCA	243
246	MetAsnLysAlaThrSerAspLysThrArgAsnVallleLysLysMetAlaLysGluGly ATGAATAAAGCTACATCTGATAAGACTCGGAATGTAATTAAAAAAATGGCAAAAGAAGGG	305
240	TACTTATTTCGATGTAGACTATTCTGAGCCTTACATTAATTTTTTTACCGTTTTCTTCCC	
306	IleTyrLeuCysValAlaGlnGlyTyrArgSerThrAlaGluGlnAsnAlaLeuTyrAla ATTTATCTTTGTGTTGCGCAAGGTTACCGCTCAACAGCGGAACAAAATGCGCTATATGCA	365
	TAAATAGAAACACAACGCGTTCCAATGGCGAGTTGTCGCCTTGTTTTACGCGATATACGT	
366	GlnGlyArgThrLysProGlyAlaIleValThrAsnAlaLysGlyGlyGlnSerAsnHis CAAGGGAGAACCAAACCTGGAGCGATTGTTACTAATGCTAAAGGTGGGCAATCTAATCAT	425
300	GTTCCCTCTTGGTTTGGACCTCGCTAACAATGATTACGATTTCCACCCGTTAGATTAGTA	-
426	AsnPheGlyValAlaValAspLeuCysLeuTyrThrSerAspGlyLysAspValIleTrp AATTTCGGTGTAGCAGTTGATTTGTGCTTGTATACGAGCGACGGAAAAGATGTTATTTGG	485
420	TTAAAGCCACATCGTCAACTAAACACGAACATATGCTCGCTGCCTTTTCTACAATAAACC	400
486	GluSerThrThrSerArgTrpLysLysValValAlaAlaMetLysAlaGluGlyPheGlu GAGTCGACAACTTCCCGGTGGAAAAAGGTTGTTGCTGCTATGAAAGCGGAAGGATTCGAA	545
400	CTCAGCTGTTGAAGGGCCACCTTTTTCCAACAACGACGATACTTTCGCCTTCCTAAGCTT	0.10
546	TrpGlyGlyAspTrpLysSerPheLysAspTyrProHisPheGluLeuCysAspAlaVal TGGGGCGGAGATTGGAAAAGTTTTAAAGACTATCCGCATTTTGAACTATGTGACGCTGTA	605
340	ACCCCGCCTCTAACCTTTTCAAAATTTCTGATAGGCGTAAAACTTGATACACTGCGACAT	003
606	SerGlyGluLysIleProThrAlaThrGlnAsnThrAsnProAsnArgHisAspGlyLys AGTGGTGAGAAAATCCCTACTGCGACACAAACACCCAATCCAAACAGACATGATGGGAAA	665
000	TCACCACTCTTTTAGGGATGACGCTGTGTTTTGTGGTTAGGTTTGTCTGTACTACCCTTT	003
666	IlevalAspSerAlaProLeuLeuProLysMetAspPheLysSerAsnProAlaArgMet ATCGTTGACAGCGCGCCACTATTGCCAAAAATGGACTTTAAATCAAATCCAGCGCGCATG	125
000	TAGCAACTGTCGCGCGGTGATAACGGTTTTTACCTGAAATTTAGTTTAGGTCGCGCGTAC	123
726	TyrLysSerGlyThrGluPheLeuValTyrGluHisAsnGlnTyrTrpTyrLysThrTyr TATAAATCAGGAACTGAGTTCTTAGTATATGAACATAATCAATATTGGTACAAGACGTAC	785
, 20	ATATTTAGTCCTTGACTCAAGAATCATATACTTGTATTAGTTATAACCATGTTCTGCATG	700

FIGURE 8 (START)

.020	TGTCGCTTGATGAAGAATTTT	
1026	ThralaAsnTyrPheLeuLys ACAGCGAACTACTTCTTAAAA	
900	ACCATGTTGTTGCCTTTTCCTATGAACCTTGAGACCATACTTTTCCTACCGACCATGATG	
966	TrpTyrAsnAsnGlyLysGlyTyrLeuGluLeuTrpTyrGluLysAspGlyTrplyrTyr TGGTACAACAACGGAAAAGGATACTTGGAACTCTGGTATGAAAAGGATGGClGGTACTAC	1025
906	TTATTGTGTTTTAACTTAAGACCCTTTTAATTTACCATACGTGGGTTATGTTTTAATCGT	-
200	AsnAsnThrLysLeuAsnSerGlyLysIleLysTrpTyrAlaProAsnThrLysLeuAla AATAACACAAAATTGAATTCTGGGAAAATTAAATGGTATGCACCCAATACAAAATTAGCA	965
846	CGTTTTCCTGCGTAGTTTCAAGCTTAATTTTCGCGCTTTCTGAATGCTTAAGGTCAAACC	903
0.46	AlalysGlyArgIlelysValArgIlelysSerAlalysAspLeuArgIleProValTrp GCAAAAGGACGCATCAAAGTTCGAATTAAAAGCGCGAAAGACTTACGAATlCCAGTTIGG	905
180	TAGTTGCTGTTTAATATGATGTACATATTCTCGAAAACGCTACAACATCGATTTTTTCTA	0.15
786	IleasnasplysleuTyrTyrMetTyrLysSerPheCysAspValValAlaLysLysAspATCAACGACAAATTATACTACATGTATAAGAGCTTTTGCGATGTTGTAGCTAAAAAAGAT	845

FIGURE 8 (END)

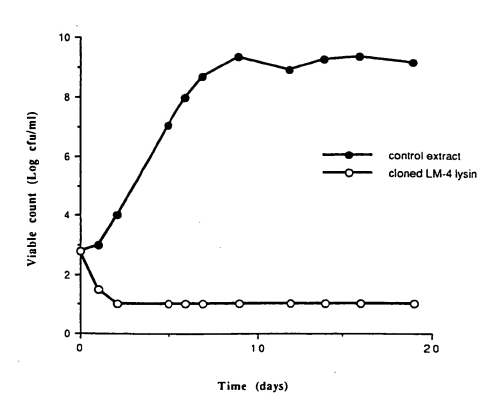


FIGURE 9

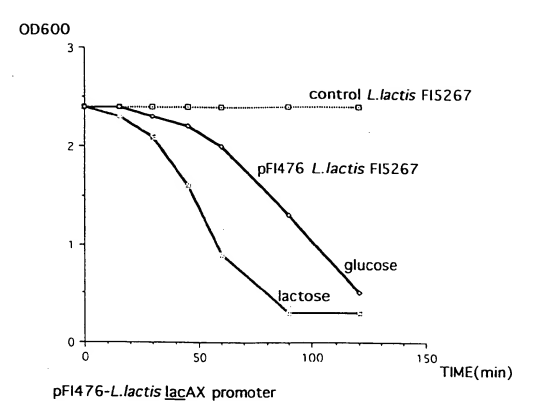
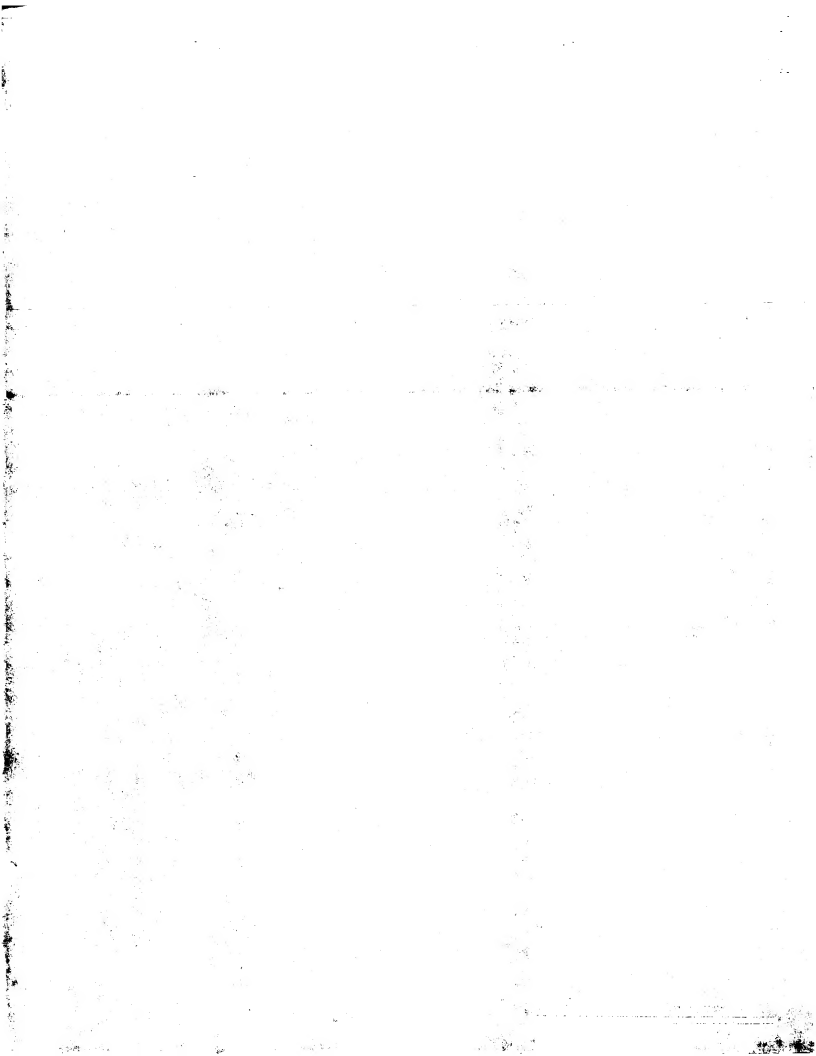


FIGURE 10



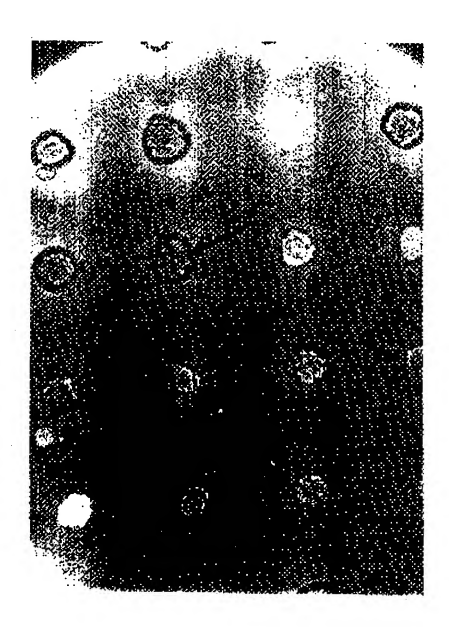


Fig. 1

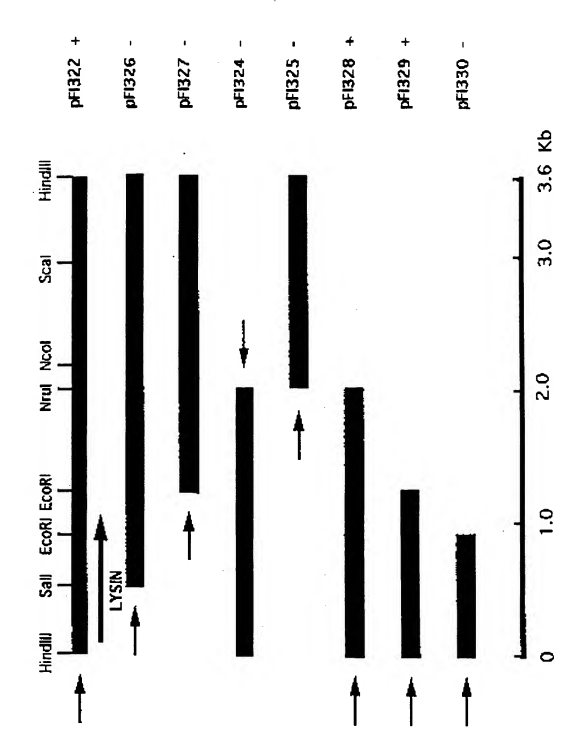


FIGURE 2

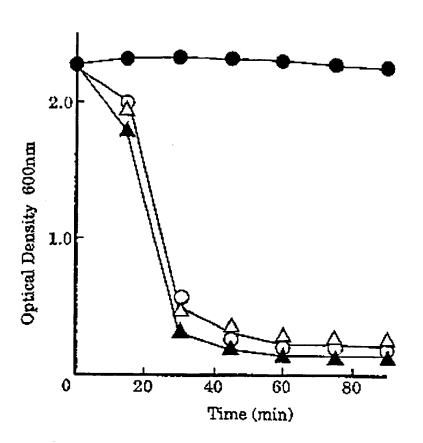


FIGURE 3

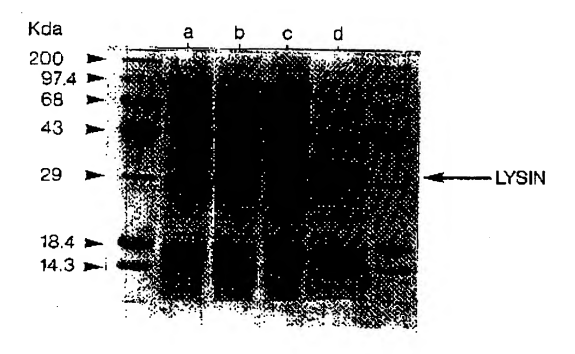


Fig. 4

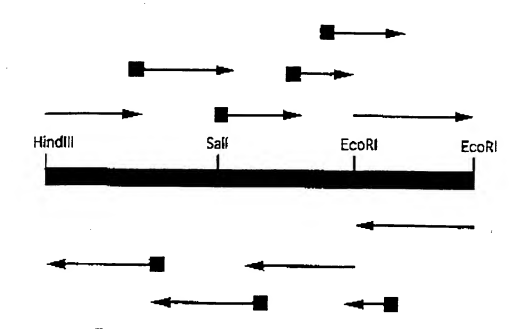


FIGURE 5

AABCTTTACT GGATAACGAC AAACTAGAAC TGACAGATGA AGGACTGAAA OBCCTTGACG AACAGCTAGC AGCATTGCAA GAAAGCGATG CTTATTTGTT TOCTCAAGAA AGCGAGGCGG GGCGGAACTG GTTTATTIGA ACAATTTACT 101 AATCGAGCTA AAAATATGG AAAGGATGAT TAATAATGGC ATTAACAGAG 151 BCATEGCTAA TTGAAAAAGC AAATCGCAAA TTGAATACET CAGGTATGAA 201 TAAABCIADA ICTBATAADA CTCGGAATGT AATTAAAAA ATGGCAAAAG 251 AAGGGATTIA TCTTTGTGTT GCGCAAGGTT ACCGET CAAC AGCGGAACAA 3D1 AATGCCCTAT ATGCACAAGD GAGAACCAAA CCTCGAGCGA ITGTTACTAA 351 TGCTAAAGGI GGGCAATCTA ATCATAATTI CGGTGTAGCA GIIGATTIGT 401 GCTTGTATAC MAGCGACGGA AAAGATGTTA TTTGGGAGTC GACAACTICC 451 CECTEGAAAA AGGITETTEC TECTAIGAAA GCGGAAGGAT TEGAAIGGGG 501 CGGAGATIGG AAAASTITIA AAGACTATCC GCATTITGAA CFAIGIGACG 551 CIGIAAGIGG IGAGAAAAIC CUTACIGCGA CACAAAACAU GAAICGAAAG 601 AGACATGATG GGAAAATCGT IGACAGCGCG CCAETATTGC CAAAAATGGA CTTFAAATCA AATCCAGCGC GCATGTATAA ATCAGGAACT GAGITCTTAG 701 TALATGAACA TAATGAATAT REGIACAAGA CETAGATGAA CEACAAATTA 751 TACTACATGE ATAGGGESS STUCCGATGET GEAGGEAAAA AAGATGCAAA AGGACGCATC ANACTTOGAN IBANANGCGC GANNGACTEN CHANTTOCAG TTTGGAATAA CACAAAATTG AATTCIGGGA AAATTAAATG GIAIGCACCC 961 AATACAAAAT TAGCAFGGTA CAACAACGGA AAAGGATAGT TGGAACICTG 951 GTATGAAAAG GAIGGCTGGT ACTACACAGC GAACIACTIC TIAAAATAAA AAGTCCCBBT TIGAGCEGGG CTTTTTATTT TCAAAGEGAG TAACAAAAAA 1051 TGTAATAAAA ATGIAATAAT CCAAGTAAGT TGTATAAAAT TIGCAGAATT AGAACGTTII ATTGATAAAT ACACCITATG AAGAGTATII TYGGCTATAT TTGCGCATTA TAGGGTTGAA TGTAACACTA TATGTAGAAT TE

RIGHER 6

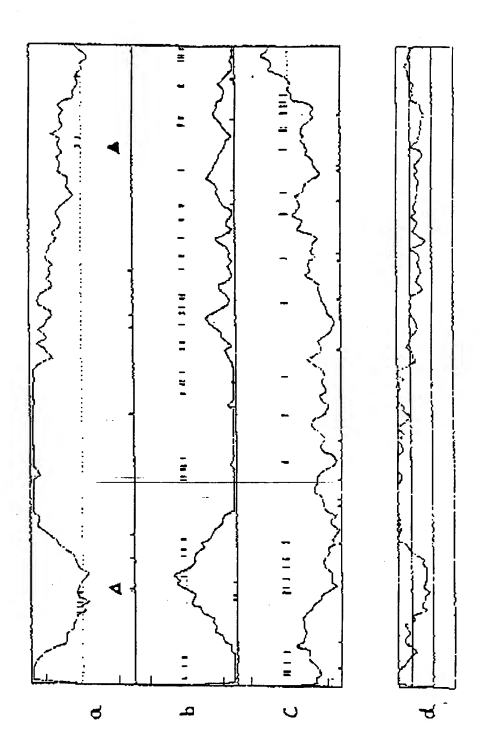


FIGURE 7

	ATGCCATTAACAGAGGCATGCCTAATTGAAAAAGCAAATCGCAAATTGAATACGTCAGGT	245
186	TACCGTAATTGTCTCCGTACCGATTAACTTTTTCGTT%AGCGTTTAACTTATGCAG1CCA	745
	MotAsnLysAlaThrSerAsptysThrArgAsnVallTcLysLysMetAlaLysGluGly ATGAATAAAGCTACAICTGATAAGACTCGGAATGTAATTAAAAAAATGGCAAAAGAAGGGG	305
246	TACTTATTTCGATGTAGACTAFTC1GAGGC1EAGATTAATTTTTTACCGTT1TCTTCCC	505
305	IletyrLeuCysValAlaGlnGlyTyrArgSerlhrAlaGluGlnAsnAlaLeuTyrAla ATTTATCTTTGIGlTGCGCAAGGITACCGCTCAACAGCGGAACAAAAIGCGCTATAJGCA	365
3U 0	TAAATAGAAACACAACGCGTT UCAA)'GGCGAGTTGTCGCCCTTGTTTTACGCGATATACGT	500
365	GlnGlyArgThrLysProGlyAfaTleValThrAsnAlaLysGlyGlyGlgGrAsnHis CAAGGGAGAACCAAACCIGGAGCGATIGTTACTAATGCTAAAGGIGGGCAALCTAATCAT	425
300	GTTCCCTCTTGGT I YGGACCTCCCTAACAATGATTACGATTTCCACCCGTYAGATTAGTA	-
	ASINPHEGRYVERARIAVORASPLBUCYSLOUTYFTRESERASPGRYLYSASPVORRER AATTTCGGTGRAGGGGTGATTTGTGCTTGTATACGAGCGACGGAAAGATGTTATTTGG	485
425	TTAAAGCCACAYCG) CAACTAAACACGAACATATGCTCGCTGCCTT11C1ACAATAAACC	70.
486	GluserThrThrSerArglrpLysLysValValAlaAlaAlaMetLysAlaGluGlyPheGlu GAGICGACAACTICCCGGIGGAAAAAGGTTGTIGCTGCTATGAAAGCGGAAGGATTGAA 	54
400	CTCAGCTG11GAAGGGCCACCTTT1TCCAACAACGACGATACT31CGCCT1CCTAAGCTT	
ć 4 E	TrpGlyGlyAsplrpLysSerPhoLysAsplyrProHisPheGluLeuCysAspAlaVal TCGGGCGGAGAIlGGAAAAGTTTTAAAGACTATCCGCATTTTGAACIAIGIGACGCTGTA	60
545	ACCCCGCCTCTAACCTTTCCAAAATTCCTGATAGGCGTAAAACETGATACACTGCGACAT	40
66¢	SerGlyGluLysIleProthrAlaIncGlnAshIhcAshProAshArgHisAspGlyLys AGTGGTGAGAAATCCCTACTGCGACACAAAACACCAATCGAAACAGACATGATGGGAAA	66
606	TCACCAGECTITTAGGGATGACGCTGIGIELLIGYGGTTAGGTTTGTCTACCACCTLI	uu
	TIEVATASOSCIATAPROLONLOUPROLYSHALASOPHHLYSSERASHPIDATAARGMCt ATCGTTGACAGCGCCCCCCTATTGCCAAAAATGGACTITAAATCAAATC	711
666	TAGCAACTG LCGCGCGGTGATAACGGTTTTTACCTGAAATTTACT1 LAGC LCCCGCGTAC	72
7.64	LyrLysSorGlyThrGluPhel euVallyrGluHisAsnGlntyrTrpTyrLysThrTyr TATAAATCAGGAACTGAGIICIIAGlaIaIGAACATAAYCAATATTGGTACAAGACGTAC	3.0
720	ATALITEAGICCTTGACTCAAGAATCATACTTGTATTAGTTATACCATGETCTCCATG	78

FIGURE 8 (START)

700	I TRASHASPLYSLBUTYFTYFMOLTYFLYSSEPPHOCYSASPYBTYBTATATACTACATGTATAGGGGCTTTTGCGATGTTGTAGCTAAAAAGAT	
786	TAGFTGCTGTTFAATATGATGTAGATATTUTCGAAAACCCCAACALCGA111111C1A	
	AlaLysGlyArgIleLysValArglleLysSerAlaLysAspl euArgIleProValTrp GCAAAAGGACGCATCAAAGTTCGAATTAAAAGCGCAAAGACTTACGAAIrccaGiilGC	
646	CRITTCCTCCGTAGTITCAAGCTTAATTITCGCGCTTTCTGAATGCTTAAGGTCAAACC	
906	ASHASHTRELYSLOUASHSDEGTYLYSTTOLYSTEDTYFAT&PFOASHTHELYSLOUATA AATAACAAAATTGAATTCTGGGAAAATTAAATGGTATGCACCCAATACAAAATTAGGA	
900	TTATTGTGTTTTAACTTAAGACCCTTTTAATTTACCATACGTGGTTATGTTTTAATCGT	
26 £	TrpTyrAsnAsnGTyLysG3yTyrLcuG1uLcuIrp1yrG3uLysAspG3y1cptyr1yr TGGTACAACAACGAAAAGGATACTTCGAACTCTGGTATGAAAAGGATGGCTGGTACTAC	
FO U	ACCATGTTGCTTTTCCTATGAACCTTGAQACCATACTTTTCCTACCGACCATGATG	
026	ThralaasmTyrPheleuLys ACAGCGAACTACTTCTTAAAA 	
J20	TGYCGCTEGATGAAGAAYTYI	

FIGURE 8 (END)

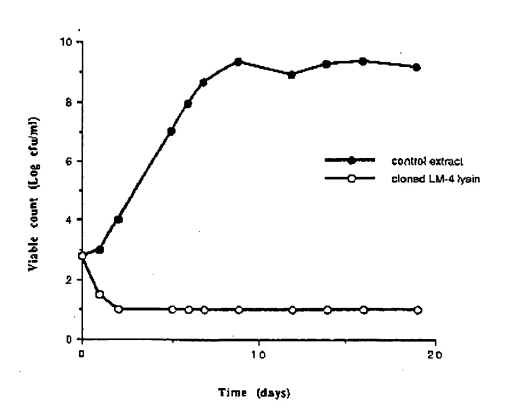


FIGURE 9

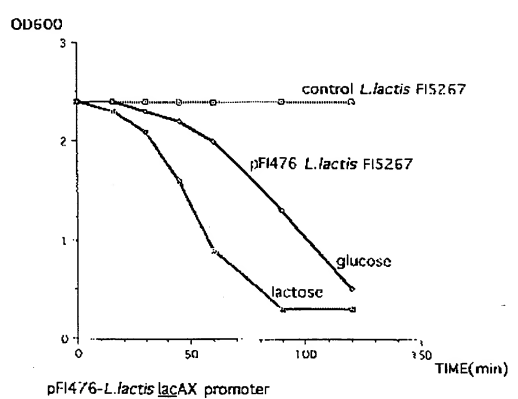


FIGURE 10

		· ·	
			•
32			
M, Y			
,			
- A ₁₁₂	-		
· ************************************			
A			
		•	
*			
i	(m m (g) + 1	gann die German von der German von d	
:			
ery	The second secon	and the state of t	S V
4			
		• ***	

.**		19	
		•	
	· ·		
	e servición de la companya de la com		
	i v		
.*	•		
		•	
		· · · · · · · · · · · · · · · · · · ·	
			*
	•		
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		